INTRODUCTION

Bioanalysis, employed for the quantitative determination of drugs and their metabolites in biological fluids, plays a significant role in the evaluation and interpretation of bioequivalence, pharmacokinetic (PK), and toxicokinetic studies. The quality of these studies, which are often used to support regulatory filings, is directly related to the quality of the underlying bioanalytical data. It is therefore important that guiding principles for the validation of these analytical methods be established and disseminated to the pharmaceutical community.

The first American Association of Pharmaceutical Scientists (AAPS)/Food and Drug Administration (FDA) Bioanalytical Workshop in 1990 focused on key issues relevant to bioanalytical methodology and provided a platform for scientific discussions and deliberations. The workshop and the report\(^1\) raised awareness of the need for validated bioanalytical methods for the regulatory acceptance of bioequivalence and pharmacokinetic data. Although the workshop addressed bioanalysis in general, it acknowledged the differences between chromatographic and ligand binding (non-chromatographic based) methods. The workshop identified the essential parameters for bioanalytical method validation, ie, accuracy, precision, selectivity, sensitivity, reproducibility, limit of detection, and stability. The outcome of the first workshop and its report resulted in improved quality of data submissions to regulatory agencies.

Following the first workshop report\(^1\) and the experience gained at the FDA, the draft Guidance on Bioanalytical Methods Validation was issued by the FDA in January 1999. This draft guidance provided stimulus and opportunity for further discussion at the 2nd AAPS/FDA Bioanalytical Workshop in January 2000. In addition, newer technology, such as chromatography coupled to tandem mass spectrometry (LC-MS/MS), was discussed along with an update on ligand-binding assays. This workshop resulted in a report “Bioanalytical Method Validation—A Revisit with a Decade of Progress”\(^2\) and formed the basis for the FDA Guidance on Bioanalytical Methods Validation in May 2001.\(^3\)

The evolution of divergent analytical technologies for conventional small molecules and macromolecules, and the growth in marketing interest in macromolecular therapies, led to the workshop held in 2000 to specifically discuss bioanalytical methods validation for macromolecules. Because of the complexity of the issues, the workshop failed to achieve a consensus. To address the need for guiding principles for the validation of bioanalytical methods for macromolecules, the AAPS Ligand-Binding Assay Bioanalytical Focus Group developed and published recommendations for the development and validation of ligand-binding assays in 2003.\(^4\)

As bioanalytical tools and techniques have continued to evolve and significant scientific and regulatory experience has been gained, the bioanalytical community has continued its critical review of the scope, applicability, and success of the presently employed bioanalytical guiding principles. The purpose of this 3rd AAPS/FDA Bioanalytical Workshop was to identify, review, and evaluate the existing practices, white papers, and articles and clarify the FDA
Guidance. The workshop addressed quantitative bioanalytical methods validation and their use in sample analysis, focusing on both chromatographic and ligand-binding assays.

GOALS AND OBJECTIVES

The purpose of the 3rd AAPS/FDA Bioanalytical Workshop was to:

• review the scope and applicability of bioanalytical principles and procedures for the quantitative analysis of samples from bioequivalence, pharmacokinetic, and comparability studies in both human and nonhuman subjects;
• review current practices for scientific excellence and regulatory compliance, suggesting clarifications and improvements where needed;
• review and evaluate validation and implementation requirements for chromatographic and ligand-based quantitative bioanalytical assays, covering all types (sizes) of molecules;
• review recent advances in technology, automation, regulatory, and scientific requirements and data archiving on the performance and reporting of quantitative bioanalytical work; and
• discuss current best approaches for the conduct of quantitative bioanalytical work regardless of the size of the molecule analyzed.

The 3rd AAPS/FDA Bioanalytical Workshop, held May 1–3, 2006, in Arlington, VA, concluded with several recommendations to achieve the above goals and objectives. While the FDA guidance remains valid, the recommendations obtained during the workshop were aimed at providing clarification and some recommendations to enhance the quality of bioanalytical work. This publication provides the clarification and recommendations obtained at the workshop with a view to achieve uniformity among the practitioners and users of quantitative bioanalysis for all types of molecules.

NONCHROMATOGRAPHIC ASSAY–SPECIFIC ISSUES

Differences Between Ligand-Binding Assays Supporting Macromolecule PK Analysis and Small Molecule Analysis by Chromatography

Ligand-binding assays (LBAs) are used throughout many organizations attempting to discover or develop new chemical entities (NCE). Besides the obvious size difference between small and macromolecule analytes, there are key structural differences. Small molecules typically are organic molecules whereas macromolecules are complex biopolymers. In addition, small molecules are prepared by organic synthesis while macromolecules are typically formed biologically. As a direct result of how macromolecules are produced, the reference standards tend to be heterogeneous, often because of posttranslational modification (eg, glycosylation or phosphorylation). In contrast, small molecule reference standards are homogeneous with a high degree of purity. Generally, small molecules are often hydrophobic and macromolecules are often hydrophilic. While chemical stability is assessed for small molecules with relative ease, macromolecule stability assessment is generally more complex, requiring the evaluation of not only chemical and physical properties, but also biological integrity (ie, is receptor binding affinity maintained?). Macromolecules are endogenous and/or structurally similar to endogenous counterparts, while small molecules are generally xenobiotics, foreign, and not present in the sample matrix. The catabolism of small molecules is typically well defined, whereas for macromolecules few specifics are known. Macromolecules typically have specific carrier proteins while small molecules can be generically bound to several endogenous proteins. Because of these significant differences between small and macromolecule analytes, different technologies, such as LC-MS for small molecules and LBAs for macromolecules, are often employed to determine drug levels for PK assessments.

Method validations for these divergent methods should consider important differences including the basis of measurement, the detection modality, and whether a sample is measured directly in the matrix or extracted before analysis. The basis of measurement of LC-MS is owed to the chemical properties of the analyte, while for LBAs, the measurement depends on a high-affinity biological binding interaction between the macromolecule analyte and another macromolecule(s) in the form of 1 or more capture/detection antibodies. Detection in LC-MS methods is direct and typically results in a linear measured response, where higher concentrations of analyte have a proportional increase in response. In contrast, the measured response in LBAs is indirect and this results in a nonlinear, often sigmoidal, measured response. Owing to the characteristics of the assay system, the calibration standard curve range for an LC-MS method is broad, often covering several orders of magnitude. In contrast, the calibration range for an LBA is typically limited to less than 2 orders of magnitude. These analyte differences, combined with the unique technologies used to measure analyte concentration, provide a strong rationale as to why consideration should be given to the need of employing some analyte-specific (small vs macromolecule) method validation guidelines.

One major point of concern in discussing the method validations for these divergent technologies centers on standards and quality control (QC) acceptance criteria (ie, the acceptable deviation from a nominal value expressed as a
Ligand-Binding Assays Pre-Study Validation

During pre-study validation, method precision and accuracy are determined through the analysis of QCs (validation samples) prepared in a biological matrix equivalent to that anticipated for study samples. Because of the endogenous nature of some biopharmaceuticals, it may be necessary to deplete the matrix of the analyte or employ a “surrogate” matrix to evaluate method accuracy and precision. One proposed validation protocol recommends that matrix be spiked at 5 or more validation sample concentrations that span the range of quantification (ie, the anticipated LLOQ, ~3 times LLOQ, mid [geometric mean], high [~75% the upper limit of quantitation, or ULOQ], and finally the anticipated ULOQ). As previously noted, the major sources of variability (imprecision and inaccuracy) differ based on technology. For LBAs, the interbatch variance component is usually a greater contributor to the overall variability than the intrabatch variance component. It is recommended that at least 2 independent determinations be made for each validation sample per assay run across a minimum of 6 independent assays runs (balance validation design). For example, 12 reportable values would result from 2 measurements across 6 independent assay runs. An appropriate statistical method should then be used to compute the summary statistics (ie, each validation sample, the repeated measurements from all runs should be analyzed together). A detailed description of this approach has been described previously.

For a method to be considered acceptable, it is recommended that both the interbatch imprecision (%CV) and the accuracy, expressed as absolute mean bias (%RE) be ±20% (25% at LLOQ and ULOQ). As an additional constraint to control method error, it is recommended that the target total error (sum of the absolute value of the %RE [accuracy] and precision [%CV] be less than ±30% ±40% at the LLOQ and ULOQ). The additional constraint of total error allows for consistency between the criteria for pre-study method validation and in-study batch acceptance. In assessing the acceptability of a method, including total error, it is not appropriate to reject assay runs. All assay runs during the validation should be included in the computation of summary statistics. The only exception would be runs rejected for cause or in cases where errors are obvious and documented.

Ligand-Binding Assays In-Study Acceptance Criteria

The recommended standard curve acceptance criteria for macromolecule LBAs are that at least 75% of the standard points should be within 20% of the nominal concentration (%RE of the back-calculated values), except at the LLOQ and ULOQ where the value should be within 25%. This requirement does not apply to “anchor calibrators,” which are typically outside the anticipated validation range of the assay and used to facilitate and improve “sigmoidal” curve-fitting.

The recommended QC acceptance criteria for macromolecule LBAs includes the use of low, medium, and high (LQC, MQC, and HQC) QCs typically run in duplicate (ie, 6 results = 3 concentrations × 2 reportable values per concentration), with assays being accepted based on a 4–6–20 rule. Exceptions to this criterion should be justified (eg, pre-study total error data approaching 30%). At least 4 of the 6 QCs must be within 20% of the nominal value. In addition, at least one QC sample per concentration needs to meet this criterion. If additional sets of QCs are used in a run, then 50% of them...
need to be “in-range” at each concentration. The following are recommendations for the placement of the controls in relation to the standard curve range. The LQC should be placed above the second nonanchor standard, ~3 times the LLOQ. The MQC is placed near the mid point (geometric not arithmetic mean) of the standard curve, while the HQC should be placed below the second nonanchor point high standard and/or ~75% of the ULOQ.

CALIBRATION CURVE AND QC RANGES
QC samples serve to monitor the performance of the methodology throughout the course of the analysis. They are the basis for demonstrating, as required in 21 CFR 320.29(a), that the analytical method is sufficiently accurate, precise, and sensitive to measure the actual concentrations achieved in the body. For studies involving pharmacokinetic profiles spanning all or most of the calibration curve, 3 QC samples run in duplicate (or at least 5% of the unknown samples), spaced across the standard curve as per the FDA Guidance, are likely sufficient to adequately monitor method performance. For an analysis where the study data fall over a small percentage of the calibration curve, it is possible that none of the QC concentrations is near the concentrations of the unknowns, thus limiting the monitoring power of the QC samples.

If a narrow range of analysis values is known or anticipated before the start of sample analysis, it is recommended that either the standard curve be narrowed and new QC concentrations used as appropriate, or if the original curve is used, existing QC concentrations be revised or sufficient QC samples at additional concentration(s) added to adequately reflect the concentrations of the study samples. Narrowing of the standard curve and preparation of new QC samples requires only a partial validation to ensure adequate performance of the new curve and QCs. A full validation is not required.

If a narrow range of analysis values is unanticipated, but observed after start of the sample analysis, it is recommended that the analysis be stopped and either the standard curve narrowed, existing QC concentrations revised, or QC samples at additional concentrations be added to the original curve before continuing with sample analysis. It is not necessary to reanalyze samples analyzed before optimizing the standard curve or QC concentrations.

CARRYOVER AND CONTAMINATION EVALUATION
Contamination, carryover, or blank response from matrix or reagents can affect the accuracy and precision of quantitation at all concentrations. However, low concentration samples are most affected as a percentage of concentration. Care should be taken to minimize interference from all contamination factors and the interference should not significantly affect the accuracy and precision of the assay.

Carryover does not necessarily involve only the next sample in the sequence. In fact, carryover from late-eluting residues on columns may affect chromatograms several samples later. Carryover from residues in rotary sampling/switching valves often appears later in the samples. Precautions should be taken to avoid contamination during sample collection and preparation. Carryover should be assessed during validation by injecting 1 or more blank samples after a high concentration sample or standard. The injector should be flushed with appropriate solvents to minimize carryover. If carryover is unavoidable for a highly retained compound, specific procedures should be provided in the method to handle known carryover. This could include injection of blanks after certain samples. Randomization of samples should be avoided, since it may interfere with the assessment of carryover problems. Contamination can be assessed by monitoring blank response in the presence of high concentration samples or standards. The assay platform (manual or automated), configuration of sampling and extraction method (e.g., manual, automated, on-line, or solid phase) in the assay should be taken into consideration when ascertaining contamination. There is no standard acceptable magnitude of carryover for a passing bioanalytical run. Carryover should be addressed in validation and minimized, and an objective determination should be made in the evaluation of analytical runs.

During validation, the operator should assess the analyte response due to blank matrix while eliminating or minimizing other contaminations. The analyte response at the LLOQ should be at least 5 times the response due to blank matrix. For immunoassays, and if the analyte is present endogenously in the matrix, the blank response can exceed 20% of LLOQ, but the contribution should not interfere with the required accuracy in the measurement of the LLOQ. In such cases, specific procedures should be provided in the method to handle blank matrix response.

DETERMINATION OF METABOLITES DURING DRUG DEVELOPMENT
A draft FDA Guidance for Industry, entitled “Safety Testing of Drug Metabolites” was issued in June 2005 by the Center for Drug Evaluation and Research (CDER). There is general support from the pharmaceutical community for the idea that a more extensive characterization of the pharmacokinetics of unique and/or major human metabolites (UMMs) would provide greater insight into the connection between metabolites and toxicological observations. This information would be best generated by the use of rugged, bioanalytical methods applied at appropriate times in drug development.
Characterization of UMMs should proceed using a flexible, “tiered” approach to bioanalytical methods validation. This tiered approach would allow metabolite screening studies to be performed in early drug development using bioanalytical methods with limited validation, with validation criteria increasing as a product moves into clinical trials. A tiered validation approach to metabolite determination would defer bioanalytical resource allocation to later in the drug-development timeline when there is a greater likelihood of drug success. As a minimum, the specifics of this tiered validation process should be driven by scientifically appropriate criteria, established a priori.

**INCURRED SAMPLE RE-ANALYSIS**

There are several situations where the performance of standards and QCs may not adequately mimic that of study samples from dosed subjects (incurred samples). Examples include metabolites converting to the parent species, protein-binding differences in patient samples, recovery issues, sample inhomogeneity, and mass spectrometric ionization matrix effects. These factors can affect both the reproducibility and accuracy of the concentration determined in incurred samples. While these effects are often characterized and minimized during method development using QC samples, it is important to ensure that they are under control when the method is applied to the analysis of incurred samples.

A proper evaluation of incurred sample reproducibility and accuracy needs to be performed on each species used for Good Laboratory Practice (GLP) toxicology experiments. It is not necessary for additional incurred sample investigations to be performed in toxicology species once the initial assessment has been performed. Incurred sample evaluations performed using samples from one study would be sufficient for all other studies using that same species.

It is generally accepted that the chance of incurred sample variability is greater in humans than in animals, so the following discussion pertains primarily to clinical studies. The final decision as to the extent and nature of the incurred sample testing is left to the analytical investigator, and should be based on an in-depth understanding of the method, the behavior of the drug, metabolites, and any concomitant medications in the matrices of interest. There should be some assessment of both reproducibility and accuracy of the reported concentration. Sufficient data should be generated to demonstrate that the current matrix produces results similar to those previously validated. It is recognized that accuracy of the result generated from incurred samples can be more difficult to assess. It requires evaluation of any additional factors besides reproducibility upon storage, which could perturb the reported concentration. These could include metabolites converted to parent during sample preparation or LC-MS/MS analysis, matrix effects from high concentrations of metabolites, or variable recovery between analyte and internal standard. If a lack of accuracy is not a result of assay performance (ie, analyte instability or interconversion) then the reason for the lack of accuracy should be investigated and its impact on the study assessed. The extent and nature of these experiments is dependent on the specific sample being addressed and should provide sufficient confidence that the concentration being reported is accurate.

The results of incurred sample reanalysis studies may be documented in the final bioanalytical or clinical report for the study, and/or as an addendum to the method validation report.

In selecting samples to be reassayed, it is encouraged that issues such as concentration, patient population, and special populations (eg, renally impaired) be considered, depending on what is known about the drug, its metabolism, and its clearance. First-in-human, proof-of-concept in patients, special population, and bioequivalence studies are examples of studies that should be considered for incurred-sample concentration verification. The study sample results obtained for establishing incurred sample reproducibility may be used for comparison purposes, and do not necessarily have to be used in calculating reported sample concentrations.

**DOCUMENTATION ISSUES**

Although the current guidance for the documentation section remains valid, further issues are now addressed and details are provided herein (Table 1) to facilitate effective documentation. Records generated during the course of method validation and study sample analysis are source records and should be retained to demonstrate the validity of the method under the conditions of use, and to support the statements made in the report. This is necessary to enable the reconstruction of the laboratory events as they occurred, since the information generated by the individual laboratory might differ from what the sponsor includes in the application.

**Documentation at the Analytical Site**

1. Documentation of standard analyte can be done by Certificate of Analysis (CA) or recertification of purity or stability data at the time of the use. In case of the internal standard, no specific CA is necessary but the lack of interference between internal standard (IS) and analyte should be established.

2. The source data (run preparation, extraction and run summary sheets, and chromatograms) of all analytical and validation runs, including failed runs should be retained.
<table>
<thead>
<tr>
<th>Items</th>
<th>Analytical Site</th>
<th>Validation Report</th>
<th>Validation Report Appendix</th>
<th>Analytical Report</th>
<th>Analytical Report Appendix</th>
</tr>
</thead>
<tbody>
<tr>
<td>Standard</td>
<td>• Certificate of analysis, purity, stability for analyte.</td>
<td>• Batch/Lot #, purity and manufacturer.</td>
<td>• Batch/Lot #, Purity and Manufacturer</td>
<td>• Stability at time of use</td>
<td>• Stability at time of use</td>
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<td></td>
<td>• Record of receipt and storage.</td>
<td>• Stability at time of use</td>
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<td></td>
<td>• Lack of interference between IS and analyte.</td>
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<tr>
<td>Stock solution preparation</td>
<td>• Records of preparation.</td>
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<tr>
<td></td>
<td>• Storage location and condition.</td>
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<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Calibrators and QCs</td>
<td>• Records of preparation.</td>
<td>• Preparation dates.</td>
<td>• SOP (optional).</td>
<td>• Short description.</td>
<td>• SOP (optional).</td>
</tr>
<tr>
<td>preparation</td>
<td>• Freezer log (sample ingress/egress, temperature).</td>
<td>• Storage conditions.</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>Run acceptance criteria</td>
<td>• SOP* for calibrators, QCs and chromatographic interferences.</td>
<td>• Short description.</td>
<td>• SOP (optional).</td>
<td>• Brief description.</td>
<td>• SOP (optional).</td>
</tr>
<tr>
<td>Assay procedure</td>
<td>• SOP for the method.</td>
<td>• Brief description of method of extraction, and analysis.</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>Sample tracking</td>
<td>• Study sample receipt, condition on receipt and location of storage.</td>
<td>• Storage condition and location.</td>
<td>• SOP (optional).</td>
<td>• Brief description.</td>
<td>• SOP (optional).</td>
</tr>
<tr>
<td></td>
<td>• Tracking of QC, calibrators and study samples.</td>
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<tr>
<td></td>
<td>• Freezer logs.</td>
<td></td>
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</tr>
<tr>
<td>Analysis</td>
<td>• Dates of extraction and analysis and instrument ID for each run.</td>
<td>• Table of runs, instrument ID, and analysis dates.</td>
<td>• Representative chromatograms.</td>
<td>• Table of all runs, and analysis dates.</td>
<td>• Chromatograms from 5%-20% of subjects for ANDA and representative chromatograms for NDA submissions.</td>
</tr>
<tr>
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<td>• Identity of QCs, calibrators, and study samples</td>
<td>• Table of calibrator results of all runs with accuracy and precision.</td>
<td>• Cross-validation, if applicable.</td>
<td>• Table of calibrator results of all passed runs with mean and % CV.</td>
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<tr>
<td></td>
<td>• Documentation of processing of calibrators, QCs, and study samples for each run</td>
<td>• Additional validation, if any.</td>
<td>• Long-term stability appended or written in a separate report.</td>
<td>• Tables of QC results of all passed runs with accuracy and precision. OK to include QC results of the failed runs.</td>
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<tr>
<td></td>
<td>• Documentation of instrument settings and maintenance.</td>
<td>• Tables of within and between run QC results (accuracy and precision).</td>
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<tr>
<td></td>
<td>• Run summary sheets.</td>
<td>• Bench-top, freeze-thaw, long-term and post-preparative and stock solution stability data.</td>
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<td>• 100% chromatograms.</td>
<td>• Extraction recovery and matrix effect.</td>
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<tr>
<td></td>
<td>• LIMS and mode of integration.</td>
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<tr>
<td></td>
<td>• Extraction dates.</td>
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Continued
Table 1. Continued

<table>
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<tr>
<th>Items</th>
<th>Analytical Site</th>
<th>Validation Report</th>
<th>Validation Report Appendix</th>
<th>Analytical Report</th>
<th>Analytical Report Appendix</th>
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<tr>
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<td>Identify runs, assay dates, and reason for failure.</td>
<td>SOP (optional).</td>
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<td>Audit trail: original and reintegration.</td>
<td>Reason for reintegration.</td>
<td>Mode of reintegration.</td>
<td>SOP (optional).</td>
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<td>Deviations from SOPs/Method</td>
<td>Documentation of deviations and unexpected events.</td>
<td>Description of deviations.</td>
<td>Impact on study results.</td>
<td>Description and supporting data of significant investigations.</td>
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<tr>
<td></td>
<td>Investigation of unexpected events.</td>
<td>Impact assessment.</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>Reassay</td>
<td>Refer to “Analysis.”</td>
<td>SOP for reassay criteria.</td>
<td>SOP (optional).</td>
<td>Table of sample IDs, reason for reassay, original and reassay values and run IDs.</td>
<td>SOP.</td>
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<tr>
<td>Communication</td>
<td>Between Analytical site and clinical site/sponsor.</td>
<td></td>
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<td></td>
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</tr>
</tbody>
</table>

SOP indicates standard operating procedures; IS, internal standard; QC, quality control; QCs, quality control samples; %CV, interbatch imprecision; ANDA, Abbreviated New Drug Application; NDA, New Drug Application, and LIMS, Laboratory Information Management Systems.

3. Reintegrated chromatograms should be explicitly identified. The reason for reintegration and the mode of reintegration should be documented. The original and reintegrated chromatograms should be retained ideally as electronic records.

4. Any problems during extraction and analysis (ie, run interruption, clogging of columns) should be identified. The appropriate remedial action should be documented.

5. In the case of multi-analyte assays (simultaneous measurement of multiple analytes in each sample), when samples are reassayed only for 1 analyte (eg, because the analyte failed to meet acceptance criteria in the original assay), the raw data collected for the other analytes should also be retained.

**Analytical/Validation Reports Should Include**

1. Summary table of all analytical runs analyzed. The tables should list the runs with run IDs, dates of analysis, whether runs passed or failed, and the reason for the failure. QC data from validation runs that only failed to meet QC acceptance criteria with no assignable cause for failure should be included in the precision and accuracy estimation.

2. Summary table of all validation runs analyzed. The tables should list the runs with run IDs, dates of analysis, whether runs passed or failed, and the reason for the failure. QC data from validation runs that only failed to meet QC acceptance criteria with no assignable cause for failure should be included in the precision and accuracy estimation.

**Source Data Documentation**

The actual conditions of use should be stated in documentation. For example, the source documentation for stability determinations during method validation should explicitly record experimental conditions such as storage temperature and duration, use of freshly prepared standard curves, and so forth. Such documentation is necessary to confirm that validation experiments support the storage conditions that existed during sample analysis.

Modification of calibration response (deletion of individual standard points that exceed predefined acceptance limits or alteration of the standard curve range) and QC levels (adding QCs or shifting in the concentration range of the study samples) should be documented with sufficient detail to...
demonstrate that the changes were justified and/or followed established procedures.

Regarding chromatographic methods, source documentation should include original and reintegrated chromatograms for accepted runs, along with the reason for changing integration parameters across a run or for individual samples within a run. Disabling electronic audit trails that record changes to integration parameters is not acceptable.

**Final Report Documentation**

A complete account of the performance of the bioanalytical method should be provided in the final report for both method validation experiments and study sample analysis. Although drug concentration data from the rejected runs need not be included in the final report, a brief description of the reasons and a tabular listing of rejected runs should be provided. The information provided would be helpful in the evaluation of the overall assay performance and acceptance of runs rejected and accepted. The final report should include a tabular listing of the actual QC results from all runs during method validation and accepted runs during study sample analysis. A table listing all reassayed samples; reason for reassy; and the values for original, reassay, and final should be included in the final report.

Currently, as described in the FDA Guidance,³ 5% to 20% of all chromatograms, including QCs samples and standards, must be submitted with an NDA or an ANDA. Because of the crucial nature of bioequivalence studies, the practice of submitting 20% of chromatograms from serially selected subjects should be continued for both NDAs and ANDAs. In general, representative chromatograms of typical analysis for other PK studies for NDAs should be sufficient for FDA submission. In circumstances where other PK studies are critical to the approval of the NDA, 20% of chromatograms may be requested for submission. However all original chromatograms and reintegrations should be retained at the site and available for audit if necessary.

Further reference is made to the FDA Guidance³ that describes the final report attributes in detail and remains generally applicable.

**STABILITY RECOMMENDATIONS**

Drug stability experiments should mimic conditions under which samples are collected, stored, and processed, as closely as possible. The experiments should be conducted in unaltered representative matrix, including the same type of anticoagulant. In cases where stripped or altered matrix is used for preparation of study calibration standards and/or QC samples, stability evaluation still must be conducted in samples prepared in unaltered matrix. If a stabilizer is normally employed with incurred samples, it should be employed with the stability samples also.

Short-term stability experiments should be designed and conducted to cover the type of storage conditions that are to be expected for study samples. This generally includes an evaluation of 3 freeze/thaw cycles, 4-hour bench top, and refrigerated stability. During freeze/thaw stability evaluations, the freezing and thawing of stability samples must mimic the intended sample-handling conditions to be used during sample analysis. If study samples are to be stored on wet ice, for thawed periods greater than 4 hours, then these conditions should be evaluated during validation as well. If during the sample analysis for a study, a sample was thawed through more than 3 cycles or if storage conditions changed and/or exceeded the sample storage conditions evaluated during method validation, stability must be established under these new conditions in order to demonstrate that the concentration values from these study samples are valid.

While short-term stability measurements are generated during method validation, long-term measurements are initiated during method validation, possibly evaluating analyte stability for a period of a few weeks, with the remaining long-term storage time points evaluated after method validation. This post-validation data can then be added to the original validation data in the form of a validation report addendum or as a stand-alone stability report. Long-term stability should be evaluated at the expected storage conditions, including expected satellite storage temperature and duration (eg, prior to shipment to the analytical laboratory). In consideration of this, there may be the need to include both −70°C and −20°C evaluations (eg, when samples are stored under different conditions at the various study locations). Refer to section entitled “Separate Stability Experiments Required at −70°C if Stability Shown at −20°C” for additional discussion.

Stability evaluations should be performed against freshly prepared standard curves. When evaluating data generated from stability experiments, intended (nominal) concentrations should be used for comparison purposes. Additionally, to determine if the initial batch of stability samples are suitable for the subsequent stability experiments, a comparison with the initial batch of stability samples is recommended. If the measured concentration of the day 0 or day 1 samples is recommended. If the measured concentration of the day 0 or 1 stability sample differs substantially from the intended concentration, this difference may be an indication that the bulk stability samples were not prepared correctly and preparing new bulk stability samples should be considered.

With respect to stock solutions prepared from certified reference standards, if the reference standard is within its expiration date when the stock solution is prepared, there is no need to prepare a new stock solution when the reference standard expires. When the stock solution exists in a different
Matrix Effects for MS-based Assays

One phenomenon influencing mass spectrometry (MS)-based bioanalytical assays is matrix effect. Matrix effect is the suppression or enhancement of ionization of analytes by the presence of matrix components in the biological samples. Quantitative measurement of matrix effect provides useful information in validation of MS-based bioanalytical methods. The quantitative measure of matrix effect can be termed as Matrix Factor (MF) and defined as a ratio of the analyte peak response in the presence of matrix ions to the analyte peak response in the absence of matrix ions, ie,

\[ \text{Matrix Factor} = \frac{\text{Peak response in presence of matrix ions}}{\text{Peak response in absence of matrix ions}} \]

An MF of 1 signifies no matrix effects. A value of less than 1 suggests ionization suppression. An MF of greater than 1 may be due to ionization enhancement and can also be caused by analyte loss in the absence of matrix during analysis. Internal standard (IS)-normalized MF is the MF of analyte divided by the MF for IS. The IS-normalized MF can also be obtained by substituting peak response with peak response ratio (analyte/IS) in the above equation for MF. Stable isotope-labeled IS minimizes the influence of matrix effects most effectively since the matrix effects observed for stable isotope-labeled IS are generally similar to those observed for the matching analyte. Analog IS may also compensate for matrix effects; however, the stable isotope-labeled internal standards are most effective and should be used whenever possible.

An absolute MF (or IS-normalized MF) of about 1 is not necessary for a reliable bioanalytical assay. However, highly variable MF in individual subjects would be a cause for the lack of reproducibility of analysis. To predict the variability of matrix effects in samples from individual subjects, determine the MF (or IS-normalized MF) for 6 individual lots of the matrix. The variability in matrix factors, as measured by the coefficient of variation should be less than 15%. If the matrix is rare and hard to obtain, the requirement for assessing variability of matrix factors in 6 lots can be waived. Stable isotope-labeled internal standards help by normalizing MF to a theoretical value of 1, and thereby reduce the effective IS-normalized MF variability. When using stable isotope IS, it is not necessary to determine the IS-normalized MF in 6 different lots.

System Suitability

Scientifically qualified and properly maintained instruments should be used for implementation of bioanalytical methods in routine drug analysis. As part of qualifying instruments, performance of system suitability ensures that the system is operating properly at the time of analysis. System suitability checks are more appropriately used for chromatographic methods to ensure that the system is sufficiently sensitive, specific, and reproducible for the current analytical run. However, the system suitability tests do not replace the required run acceptance criteria with calibration standards and QC samples. System suitability tests, when appropriate, are recommended to ensure success, but are not required, nor do they replace the usual run acceptance criteria.

Reference Standards

Analytical reference standards are used for the preparation of calibration standards and QC samples. Reference standard lot numbers, purity, storage, stability, handling, and supporting documentation should be monitored and maintained. Reference standards should be used before their expiration or recertification dates. Some compounds used as
internal standards or rare metabolites are available in very small amounts, and their certificates of analysis (CA) may not be available. If the full CA is not available for rare metabolites, at a minimum, the documented purity information should be obtained. CA or purity information of internal standards is not always necessary for the use of internal standards. When purity information is not available for the internal standard, it needs to be demonstrated that the internal standard does not interfere with the chromatography of the analyte(s) of interest. Macromolecular reference standards are often heterogeneous and may present unique comparability and stability considerations.

**Cross-Validation Required When Moving a Method Between LC-MS/MS Instruments**

Moving from different models of instrument (eg, Sciex API-4000 to a Sciex API-5000) would require cross-validation, but there was very little support for requiring cross-validation when switching between equivalent instruments. It was noted that a qualification experiment is usually performed by most analysts before samples are run on a given instrument and that these experiments were usually sufficient to allow the qualification of a new instrument for a specific assay.

**Specific Criteria for Cross-Validation**

The term cross-validation was used liberally throughout the 3rd AAPS/FDA Bioanalytical Workshop when considering all different types of changes in bioanalytical methods. Cross-validation was discussed when using matrix from different species, when using matrix with different anticoagulants, when transferring methods to other laboratories and when transferring methods to other analysts. However no specific strategies, procedures, and acceptance criteria were discussed to adequately perform these cross-validation experiments. Cross-validation procedures and acceptance criteria need to remain flexible, considering the various bioanalytical situations where it would be required. Specific cross-validation criteria should be established a priori via a standard operating procedure.

**Separate Stability Experiments Required At −70°C if Stability Shown at −20°C**

A significant group of workshop attendees believed that biological sample stability shown at a given temperature (eg, about −20°C), automatically defined sample stability at a lower temperature (eg, about −70°C) based on Arrhenius principles of chemical reactivity. These principles dictate lower rates of chemical reactivity (ie, analyte degradation) occurring at lower temperatures. The argument against assuming sample stability at lower temperatures was based on matrix degradation rather than chemical stability of the analyte. An effective (if anecdotal) argument was made during the meeting, which questioned the stability of biological matrix proteins at lower temperatures. It was argued that a lower temperature could cause denaturation or precipitation of matrix proteins and that this could affect protein binding or the ability to extract the drug from the matrix. There was general agreement that sample matrix will often have a different consistency depending on freezing conditions. Based on these arguments, there did seem to be some support for validating stability at lower storage temperatures even if stability has already been determined at a higher temperature. Additional stability at lower temperature should be

**RUN ACCEPTANCE CRITERIA**

The elements of chromatographic run acceptance previously described in the FDA Guidance3 are listed in Table 2. Recommendations specific to LBAs are also provided in the section “Ligand-Binding Assays In-Study Acceptance Criteria.”

**VALIDATION TOPICS WITH NO CONSENSUS**

This section describes the topics for which no consensus could be reached during the 3rd AAPS/FDA Bioanalytical Workshop. The purpose of this section is to provide comments and emphasize that further discussion and direction will be necessary.

**Cross-Validation Of Bioanalytical Methods When Using Different Anticoagulant Counter-Ions**

There was recognition of a distinct difference between EDTA and heparin-containing plasma and that a bioanalytical method validated for one could not be used for the other without some revalidation of the method but no consensus was reached for the need for cross-validation when using the same anticoagulant with a different counter-ion. As an example, attendees could not agree on the degree of cross-validation necessary for a method validated using sodium-heparinized plasma when it was applied to a lithium-heparinized sample.

**Cross-Validation Required When Using Different Strains or Sexes of a Species**

No decisive arguments came forward in support of this activity although it was agreed that there could be some differences in biological matrix originating from the different strains or sexes. The general trend of the debate was that validation experiments to address such differences should not be considered the norm and should be performed when there are method-related concerns that can be attributed to a specific strain or sex-related difference.
### Table 2. Routine Drug Analysis Process and Run Acceptance Criteria

<table>
<thead>
<tr>
<th>Process or Criteria</th>
<th>Chromatographic Assays</th>
<th>Ligand-Binding Assays</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Preparation of standards and QC samples</td>
<td>Standards and QC samples can be prepared from the same spiking stock solution, provided the solution stability and accuracy have been verified. A single source of matrix may also be used, provided selectivity has been verified.</td>
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<tr>
<td>2. Placement of samples</td>
<td>Standard curve samples, blanks, QCs, and study samples can be arranged as considered appropriate within the run, and support detection of assay drift over the run.</td>
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<tr>
<td>3. Number of calibration standards in a run</td>
<td>Include with each analytical batch:</td>
<td>Include with each analytical batch or micro-titer plate:</td>
</tr>
<tr>
<td></td>
<td>• Blank matrix (sample without internal standard)</td>
<td>• Blank matrix</td>
</tr>
<tr>
<td></td>
<td>• Zero standard (matrix sample with internal standard)</td>
<td>• Non-zero calibration standards: A minimum of 6 standard points. Can include anchor points (below LLOQ or above ULOQ in the asymptotic low- and high- concentration end of the standard curve)</td>
</tr>
<tr>
<td></td>
<td>• Non-zero calibration standards: A minimum of 6 standard points</td>
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<tr>
<td>4. Acceptance criteria for calibration standards</td>
<td>Residuals (absolute difference between the back calculated and nominal concentration) for each calibration standard should meet the following limits:  • LLOQ standard &lt; 20%  • All other standards &lt; 15%  A minimum of 75% standards (at least 6 nonzero points) should be within the above limits for the analytical run to qualify. Values falling outside these limits can be discarded, provided they do not change the established model.</td>
<td>Residuals for each calibration standard should meet the following limits:  • LLOQ and ULOQ standards &lt; 25%  • All other standards &lt; 20%  • Any anchor points if used, are not to be included in the above acceptance criteria.</td>
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<tr>
<td>5. Number of QC samples in a batch</td>
<td>Include QC samples at the following 3 concentrations (within the calibration range) in duplicate with each analytical batch:  • Low: near the LLOQ (up to 3× LLOQ)  • Medium: midrange of calibration curve  • High: near the high end of range  Each analytical batch should contain 6 or a minimum of 5% of the total number of unknown samples. Add QCs in multiples of three concentrations (low, medium, high) when needed.</td>
<td>QC samples at the following 3 concentrations (within the calibration range) in duplicate should be added to each microtiter plate:  • Low: above the second nonanchor standard, ~ 3× LLOQ  • Medium: midrange of calibration curve  • High: below the second nonanchor point high standard at ~75% of ULOQ</td>
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<tr>
<td>6. Acceptance criteria for QC samples*</td>
<td>Allowed % deviation from nominal values:  • QCs prepared at all concentrations greater than LLOQ &lt; 15%  • Low QC (if prepared at LLOQ) &lt; 20%  At least 67% (4 of 6) of the QC samples should be within the above limits; 33% of the QC samples (not all replicates at the same concentration) can be outside the limits. If there are more than 2 QC samples at a concentration, then 50% of QC samples at each concentration should pass the above limits of deviation.</td>
<td>Allowed % deviation from nominal values:  • QCs prepared at all concentrations other than LLOQ and ULOQ &lt; 20%  • Low and high QC (if prepared at LLOQ or ULOQ) &lt; 25%  In certain situations wider acceptance criteria may be justified, eg, when total error during assay validation approaches 30%</td>
</tr>
</tbody>
</table>

Continued

Table 2. Continued

<table>
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<tbody>
<tr>
<td>7. Replicate analysis</td>
<td>In general, samples can be analyzed with a single determination without replicate analysis if the assay method has acceptable variability as defined by the validation data. Duplicate or replicate analysis can be performed for a difficult procedure where high precision and accuracy may be difficult to obtain.</td>
<td>Accuracy can generally be improved by replicate analysis. Therefore, duplicate analysis is recommended. If replicate analysis is performed, the same procedure should be used for samples and standards.</td>
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<tr>
<td>8. Multiple analytes in a run†</td>
<td>Samples involving multiple analytes in a run should not be rejected based on the data from 1 analyte failing the acceptance criteria.</td>
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<tr>
<td>9. Rejected runs</td>
<td>The data from rejected runs need not be documented, but the fact that a run was rejected and the reason for failure should be reported.</td>
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</table>

*The issue of monitoring the effect of sample dilution is referred to in the FDA guidance.† The guidance indicates that if the dilutions are conducted with like matrix (human plasma for human plasma), no within-study dilution matrix QC samples are necessary. However, the extent to which samples are allowed to be diluted should be tested during validation. If tested during validation, there is no need to run dilution QCs up to the tested dilution factor during sample analysis. If during sample analysis it is determined that the required dilution factor is greater than the extent tested during validation, an additional dilution factor should be tested during sample analysis. On the other hand, if the dilution is allowed and performed with an unlike matrix, QC samples should be diluted in the same manner as the study samples, and should be analyzed with the diluted samples. All diluted QCs should be created within the assay calibration range, and similar acceptance criteria as defined here should be used, unless alternate specific criteria can be justified.

† Multiple analyte assessment in a single analytical run was a topic discussed at the meeting. Although the FDA Guidance indicates that a run should not be rejected for the remaining analytes if one fails, it does not address how to assess and report all analyte concentrations upon reanalysis of the failed analyte(s). In this regard, concentrations from the first accepted run should be reported and if this analyte is repeated in simultaneous assays when analyzing for different analytes, it is not necessary to quantitate the already reported analytes. However, the source data from all acceptable runs, regardless of whether the concentrations from these runs were reported or not, should be retained.

QC indicates quality control; QCs, quality control sample; LLOQ, lower limit of quantitation and ULOQ, upper limit of quantitation.

required for macromolecules and may also be performed for small molecules as needed.

**Stability Criteria for Stock Solution Stability**

The need to characterize the stability of stock solutions was emphasized throughout the meeting and accepted as a core validation experiment. However, there was no agreement on the degree of degradation that defines acceptable stability. The consensus was that lower degradation in the standard acceptable ranges is desirable since these stock solutions are used for making other solutions and this error may be propagated in the concentrations reported for biological samples.

**Acceptance Criteria for Internal Standards**

The practice of placing precision criteria on internal standards as an additional run acceptance test was discussed. A highly variable internal standard can be an indication of an uncontrolled process during sample analysis, especially if the internal standard response is variable with incurred samples. It is recognized that the internal standard is present in a bioanalytical assay to compensate for variability of extraction in LC-MS/MS analysis. This is most likely to be the case when the internal standard is isotopically labeled. When assessing the impact of internal standard variability, it is important to determine, in cases of low internal standard response, that the assay continues to have the ability to accurately quantify at the LLOQ. No agreement was reached on the inclusion of internal standard criteria or on the magnitude of acceptable internal standard precision. However, if study samples or analytical runs are rejected or repeated based on internal standard response variability, objective criteria are necessary and need to be established a priori.

**CONCLUSION**

For quantitative bioanalytical method validation procedure and requirements, there was a relatively good agreement between chromatographic assays and ligand-binding assays. It was realized that the quantitative and qualitative aspects of bioanalytical method validation should be reviewed and applied appropriately.

1. Some of the major concerns between the 2 methodologies related to the acceptable total error for
precision and accuracy determination and acceptance criteria for an analytical run. The acceptable total error for precision and accuracy for both the methodologies is less than 30. The 4-6-15 rule for accepting an analytical run by a chromatographic method remained acceptable while a 4-6-20 rule was recommended for ligand-binding methodology.

2. The 3rd AAPS/FDA Bioanalytical Workshop clarified the issues related to placement of QC samples, determination of matrix effect, stability considerations, use of internal standards, and system suitability tests.

3. There was a major concern and issues raised with respect to stability and reproducibility of incurred samples. This should be addressed for all analytical methods employed. It was left to the investigators to use their scientific judgment to address the issue.

4. In general, the 3rd AAPS/FDA Bioanalytical Workshop provided a forum to discuss and clarify regulatory concerns regarding bioanalytical method validation issues.

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*The views expressed in this article are those of the authors and do not reflect official policy at the FDA. No official endorsement by the FDA is intended or should be inferred.

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